
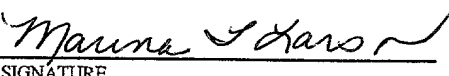


FORM PTO-1390 (REV. 11-2000)		U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY'S DOCKET NUMBER SELF. P-016-3
TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371			U.S. APPLICATION NO (if known, see 37 CFR 1.5) 09/869887
INTERNATIONAL APPLICATION NO. PCT/US00/00620	INTERNATIONAL FILING DATE 11 January 2000	PRIORITY DATE CLAIMED 01 December 1999	
TITLE OF INVENTION Disposable Test Strips With Integrated Reagent/Blood Separation Layer			
APPLICANT(S) FOR DO/EO/US McAleer, et al			
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:			
<ol style="list-style-type: none"> 1. <input checked="" type="checkbox"/> This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2. <input type="checkbox"/> This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. 3. <input type="checkbox"/> This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (21) indicated below. 4. <input type="checkbox"/> The US has been elected by the expiration of 19 months from the priority date (Article 31). 5. <input checked="" type="checkbox"/> A copy of the International Application as filed (35 U.S.C. 371(c)(2)) <ol style="list-style-type: none"> a. <input type="checkbox"/> is attached hereto (required only if not communicated by the International Bureau). b. <input type="checkbox"/> has been communicated by the International Bureau. c. <input checked="" type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US). 6. <input type="checkbox"/> An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)). <ol style="list-style-type: none"> a. <input type="checkbox"/> is attached hereto. b. <input type="checkbox"/> has been previously submitted under 35 U.S.C. 154(d)(4). 7. <input checked="" type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)) <ol style="list-style-type: none"> a. <input type="checkbox"/> are attached hereto (required only if not communicated by the International Bureau). b. <input type="checkbox"/> have been communicated by the International Bureau. c. <input type="checkbox"/> have not been made; however, the time limit for making such amendments has NOT expired. d. <input checked="" type="checkbox"/> have not been made and will not be made. 8. <input type="checkbox"/> An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). 9. <input type="checkbox"/> An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10. <input type="checkbox"/> An English language translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). 			
Items 11 to 20 below concern document(s) or information included:			
<ol style="list-style-type: none"> 11. <input type="checkbox"/> An Information Disclosure Statement under 37 CFR 1.97 and 1.98. 12. <input type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 13. <input checked="" type="checkbox"/> A FIRST preliminary amendment. 14. <input type="checkbox"/> A SECOND or SUBSEQUENT preliminary amendment. 15. <input type="checkbox"/> A substitute specification. 16. <input type="checkbox"/> A change of power of attorney and/or address letter. 17. <input type="checkbox"/> A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821 - 1.825. 18. <input type="checkbox"/> A second copy of the published international application under 35 U.S.C. 154(d)(4). 19. <input type="checkbox"/> A second copy of the English language translation of the international application under 35 U.S.C. 154(d)(4). 20. <input type="checkbox"/> Other items or information: 			

U.S. APPLICATION NO. 09/869887 INTERNATIONAL APPLICATION NO. PCT/US00/00620		ATTORNEY'S DOCKET NUMBER SELF.P-016-3	
21. <input checked="" type="checkbox"/> The following fees are submitted: BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)): Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO \$1000.00 International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO \$860.00 International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO \$710.00 International preliminary examination fee (37 CFR 1.482) paid to USPTO but all claims did not satisfy provisions of PCT Article 33(1)-(4) \$690.00 International preliminary examination fee (37 CFR 1.482) paid to USPTO and all claims satisfied provisions of PCT Article 33(1)-(4) \$100.00 ENTER APPROPRIATE BASIC FEE AMOUNT =		CALCULATIONS PTO USE ONLY <div style="border: 1px solid black; padding: 5px; width: fit-content; margin: 0 auto;">\$ 100⁰⁰</div>	
Surcharge of \$130.00 for furnishing the oath or declaration later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(e)).		\$	
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE
Total claims	10 - 20 =	0	x \$18.00
Independent claims	1 - 3 =	0	x \$80.00
MULTIPLE DEPENDENT CLAIM(S) (if applicable)		+ \$270.00	
TOTAL OF ABOVE CALCULATIONS =		\$	
<input type="checkbox"/> Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.		+	
SUBTOTAL =		\$	
Processing fee of \$130.00 for furnishing the English translation later than <input type="checkbox"/> 20 <input type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(f)).		\$	
TOTAL NATIONAL FEE =		\$	
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property +		\$	
TOTAL FEES ENCLOSED =		\$ 100 ⁰⁰	
		Amount to be refunded:	\$
		charged:	\$
a. <input checked="" type="checkbox"/> A check in the amount of \$ <u>100.00</u> to cover the above fees is enclosed. b. <input type="checkbox"/> Please charge my Deposit Account No. _____ in the amount of \$ _____ to cover the above fees. A duplicate copy of this sheet is enclosed. c. <input type="checkbox"/> The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. <u>15-0610</u> A duplicate copy of this sheet is enclosed. d. <input type="checkbox"/> Fees are to be charged to a credit card. WARNING: Information on this form may become public. Credit card information should not be included on this form. Provide credit card information and authorization on PTO-2038.			
NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137 (a) or (b)) must be filed and granted to restore the application to pending status.			
SEND ALL CORRESPONDENCE TO: OPPEDahl & LARSON LLO  021121 PATENT TRADEMARK OFFICE		<div style="text-align: center;">  SIGNATURE Marina T. Larson, Ph.D. NAME 32,038 REGISTRATION NUMBER </div>	

09/869887

JUL 05 2001

SELF.P-016-3
PATENT APPLICATION

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant : McAleer, et al.
Serial No. : TBA
Filing Date: : Herewith
Title : Disposable Test Strips With Integrated Reagent/Blood
Separation Layer

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents
U.S. Patent and Trademark Office
Washington, D.C. 20231

Sir:

This application is based upon the Claims as amended during the Preliminary Examination.
A copy of said examination is attached for the Examiner's convenience.

Preliminary to the calculation of fees and the examination, please cancel claims 11 through
23.

Respectfully submitted,

OPPEDAHL & LARSON LLP

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I hereby certify that this paper and the attachments named herein are being deposited with the United States Postal Service as
Express Mail in an envelope addressed to Commissioner of Patents, U.S. Office of Patents and Trademarks, Washington, D.C.
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7/5/01
Date of Signature

Linda L. Orr
Linda L. Orr

EL556130787US

DISPOSABLE TEST STRIPS WITH INTEGRATED
REAGENT/BLOOD SEPARATION LAYER

DESCRIPTION

BACKGROUND OF THE INVENTION

This application relates to disposable test strips for use in electrochemical determinations of blood analytes such as glucose, and to methods and compositions for use in making such strips.

5 Glucose monitoring is a fact of everyday life for diabetic individuals, and the accuracy of such monitoring can literally mean the difference between life and death. To accommodate a normal life style to the need for frequent monitoring of glucose levels, a number of glucose meters are now available which permit the individual to test the glucose level in a small amount of blood.

10 Many of these meters detect glucose in a blood sample electrochemically, by detecting the oxidation of blood glucose using an enzyme such as glucose oxidase provided as part of a disposable, single-use electrode system. Examples of devices of this type are disclosed in European Patent No. 0 127 958, and US Patents Nos. 5,141,868, 5,286,362, 5,288,636, and 5,437,999 which are
15 incorporated herein by reference for purposes of those countries which permit such incorporation.

20 In general, existing glucose test strips for use in electrochemical meters comprise a substrate, working and reference electrodes formed on the surface of the substrate, and a means for making connection between the electrodes and the meter. The working electrode is coated with an enzyme capable of oxidizing glucose, and a mediator compound which transfers electrons from the enzyme to the electrode resulting in a measurable current when glucose is present. Representative mediator
25 compounds include ferricyanide, metallocene compounds such as ferrocene, quinones, phenazinium salts, redox indicator DCPIP, and imidazole-substituted osmium compounds.

Working electrodes of this type have been formulated in a number of ways. For example, mixtures of conductive carbon, glucose oxidase and a mediator

have been formulated into a paste or ink and applied to a substrate. EP 0 127 958 and US 5,286,362. In the case of disposable glucose strips, this application is done by screen printing in order to obtain the thin layers suitable for a small flat test strip. The use of screen printing, however, introduces problems to the operation of the electrode.

5 Unlike a thicker carbon paste electrode which remains fairly intact during the measurement, screen printed electrodes formed from carbon pastes or inks are prone to break up on contact with the sample. The consequences of this breakup are two-fold. Firstly, the components of the electrode formulation are released into solution. Once these components drift more than a diffusion length away from the underlying conductive layer, they no longer contribute toward the measurement, but in fact diminish the response by depleting inwardly-diffusing analyte. Secondly, the breakup of the screen printed electrode means that the effective electrode area is falling over time.

10 The combination of these two effects results in current transients which fall rapidly from an initial peak over the period of the measurement, and a high sensitivity to oxygen which quickly competes with the mediator for the enzyme. This fact is clearly demonstrated by the much lower currents measured in blood samples than in plasma samples or other aqueous media, and can result in erroneous readings. A further consequence is that the transients are often "lumpy" as the electrode breaks up in a chaotic manner. Lumpy transients either give rise to erroneous readings or rejected strips, neither of which are acceptable.

15 In addition to the potential for electrode breakup of screen-printed carbon-based electrodes, known electrodes used in disposable glucose test strips have been kinetically-controlled, i.e., the current depends on the rate of conversion of glucose by the enzyme. Because the response measured by the instrument represents a balance between the reactions of enzyme and mediator, enzyme and glucose and enzyme and oxygen, and because each of these reactions has its own dependence on temperature, the response of a kinetically-controlled test strip is very sensitive to the temperature of the sample. Substantial variation in the measured glucose value can therefore occur as a result of variations in sample handling.

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A further challenge facing sensors for electrochemical glucose detection arises as a result of interference from blood cells present in the sample. The level of red blood cells is reflected in the hematocrit reading. Typically, high hematocrit samples results in readings that are lower than the true value, while low hematocrit samples result in readings that are higher because the blood cells tend to foul the surface of the electrode and limit electron transfer. Also, oxygen bound to the hemoglobin of red blood cells competes with the mediator for the reduced enzyme, thereby further diminishing the glucose response. Attempts have been made to limit the hematocrit effect by adding a membrane to filter out blood components (see, US Patent No. 5,658,444, which is incorporated herein by reference for purposes of those countries which permit such incorporation), but this adds an extra step to the manufacturing process, with associated increase in cost and often degraded performance in other areas such as precision.

Because of the importance of obtaining accurate glucose readings to the well-being of a patient using the meter and disposable test strips, it would be highly desirable to have a glucose test strip which did not suffer from these drawbacks, and which therefore provided a more consistent and reliable indication of actual blood glucose values, regardless of actual conditions. It is therefore an object of the present invention to provide disposable glucose test strips which provide a glucose reading that is essentially independent of the hematocrit of the sample, and which include an integrated reagent/blood separation layer.

It is a further object of the present invention to provide an improved method for making disposable glucose test strips.

SUMMARY OF THE INVENTION

The present invention provides an improved disposable test strip for use in a test meter of the type which receives a disposable test strip and a sample of blood from a patient and performs an electrochemical analysis of the amount of a blood analyte such as glucose in the sample. The test strip comprises:

- (a) a substrate;
- (b) a first conductive element disposed on the substrate;

(c) a second conductive element disposed on the substrate in sufficient proximity to the first conductive element to allow the completion of an electrical circuit between the first and second conductive elements when a sample of blood is placed on the test strip;

(d) a non-conductive integrated reagent/blood separation layer disposed over the first conductive element; and

(e) contacts for making an electrical connection between the first and second conductive elements and the test meter. The integrated reagent/blood separation layer comprises reagents for the electrochemical detection of the analyte dispersed in a non-conductive matrix effective to exclude blood cells from the surface of the first conductive element while permitting access to the first conductive element by soluble electroactive species. In one embodiment of the invention, a glucose test strip is formed with an integrated reagent/blood separation layer comprising a filler which has both hydrophobic and hydrophilic surface regions, an enzyme effective to oxidize glucose, e.g., glucose oxidase, and a mediator effective to transfer electrons from the enzyme to the conductive element. The filler is selected to have a balance of hydrophobicity and hydrophilicity such that on drying the integrated reagent/blood separation layer forms a two-dimensional network on the surface of the conductive element. Preferred integrated reagent/blood separation layers comprise non-conductive silica fillers in combination with materials such as hydroxyethyl cellulose (HEC). The silica and HEC form a two-dimensional network which excludes red blood cells, thus rendering the test strip substantially insensitive to the hematocrit of the patient.

In a preferred embodiment of the invention, the test strips are prepared with an insulation layer disposed over at least the first conductive element. This insulation layer has an aperture formed in it which is aligned with a portion of the first conductive element, and the integrated reagent/blood separation layer is disposed to make contact with the first conductive element through this aperture.

BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A and 1B show an electrode structure useful in a disposable test strip in accordance with the invention;

Fig. 2 shows a test strip in accordance with the invention;

Figs. 3A - 3C show the current measured as a function of glucose concentration for three different hematocrit levels;

Fig. 4 shows the relationship of the glucose-concentration dependence of the measured current as a function of hematocrit;

Figs. 5A - 5C show the current measured as a function of glucose in blood and a control solution for three different conductive elements;

Figs. 6A and 6B show the current measured as a function of glucose at two different temperatures;

Fig. 7 shows a further embodiment of a glucose test strip according to the invention;

Figs 8A and 8B show current transients observed using a test strip according to the invention and a commercial carbon-based test strip;

Figs. 9A-C show a three-step process for manufacture of test strips in accordance with the invention; and

Figs. 10A-10G show the manufacture of a test strip in accordance with the invention.

DETAILED DESCRIPTION OF THE INVENTION

Figs. 1A and 1B show electrodes useful in a disposable test strip in accordance with the invention. As shown, the electrodes are formed on a substrate 10. On the substrate 10 are placed two conductive elements 14' and 16, connected by leads 14 and 15 to conductive contacts 11, 12, and 13. An insulating mask 18 is then formed, leaving at least a portion of conductive elements 14' and 16, and the contacts 11, 12 and 13 exposed. A non-conductive integrated reagent/blood separation layer 17 is then applied over the insulating mask 18 to make contact with conductive element 16.

The assembly shown in Fig. 1 provides a fully functional assembly for the measurement of a blood analyte when connected to a meter. Advantageously, however, the electrode strips of the invention are finished by applying a nylon or polyester mesh 21 over the sample application region defined by the location of the integrated reagent/blood separation layer 17 of the electrode assembly 22, and then a top cover 23 to prevent splashing of the blood sample. (Fig. 2) The polyester mesh acts to guide the sample to the reference electrode, conductive element 14', thereby triggering the device and initiating the test.

The utilization of a non-conductive integrated reagent/blood separation layer provides an important distinction from and advantage over known test strips which utilize a conductive reagent-containing slurry to print the reagents. In these systems, the printed slurry becomes a functional part of the electrode and charge transfer can take place at the outer surface of the reagent layer. If the layer is in direct contact with blood, i.e., when no intervening separation layer has been deposited, red and white blood cells, fat and proteins present in the sample can interact with the reagent layer and interfere with the measurement of the amount of analyte in the sample.

In contrast, in the present invention, the integrated reagent/blood separation layer is non-conductive, and thus is not a part of the electrode either structurally or functionally. Charge transfer does not occur unless electroactive species pass through the openings/pores of the integrated reagent/blood separation layer to reach the underlying conductive element. Thus, the integrated reagent/blood separation layer provides a barrier to the passage of interferents such as cells and macromolecules to the conductive element resulting in a device with superior properties that is simpler to make.

In achieving this result, it is particularly desirable that the integrated reagent/blood separation layer be deposited in such a way that no portion of the conductive element 16 be directly exposed to the sample when it is placed in the sample application region. The methodology described above, in which an insulating layer with apertures providing access to the conductive elements 14' and 16 is utilized is particularly suited for achieving this result. Thus, as shown in Figs. 9A-C, this

methodology allows the formation of the test strip in only three steps. In the first step (Fig. 9A), two conductive elements 14' and 16 and associated leads and contacts are deposited on a substrate. In a second step (Fig. 9B), a layer of insulating material is deposited over the conductive elements. The insulating material has two apertures 94 and 96, one in alignment with each of the conductive elements 14' and 16. In the third step, (Fig. 9C), the integrated reagent/blood separation layer 17 is deposited over the aperture 96. By making the deposited layer 17 larger in dimensions than the aperture 96, the reagent layer completely covers the underlying conductive element such that it is not exposed directly to the sample, thereby providing effective blood separation.

The complete coverage of conductive element 16 also addresses another source of error which can occur as a result of electrochemical oxidation or small molecules such as ascorbic acid, uric acid and acetaminophen which may be present in the sample. When present, the oxidation of these molecules at the surface of the electrode leads to spuriously elevated current levels, and thus an inaccurate measurement of the desired analyte, e.g. glucose. The integrated reagent/blood separation layer of invention will not generally exclude these molecules, since they are small compared to the pore sizes observed. However, by including a pH buffer in the integrated reagent/blood separation layer one can shift the local pH at the electrode surface to a level where electrochemical potential of these species is higher. Thus, for example, the use of an integrated reagent/blood separation layer in which the pH is buffered to a level of around pH 5 will substantially reduce the impact of these interferents. To maximize the effectiveness of this buffering, however, the entire conductive element must be covered, since even a relatively small region of exposed (not buffered) electrode surface can result in a large interference current.

Not only do the test strips of the invention provide performance benefits resulting from the separation of the conductive element from the blood sample, the test strips of the invention are also resistant to other sources of error. For example, during the period of a test, reagents may diffuse laterally away from the original deposit. If the reagent layer is deposited directly on the conductive element, these reagents will continue to contribute to the measured signal. Any variations in

convective diffusion from test to test (for example as a result of differences in temperature or differences in the handling of the instrument) will therefore be manifested as irreproducibility in the signal. If the reagent layer overlaps the insulation print, however, lateral diffusion away from the aperture will not contribute to the signal and therefore will not give rise to variations in the signal.

In addition to providing a test strip with beneficial properties, the methodology outlined in Figs. 9A-C offers several advantages from a manufacturing perspective. First, if the reagent layer is printed directly onto the conductive element, the "active area" is defined by the area of the reagent layer. The precision of the test is therefore determined by the precision with which the reagent layer can be printed. In contrast, by first depositing an apertured insulation layer defining the region of the contact between the reagent layer and the underlying conductive element, the active area is defined by the size of the aperture in the insulation layer. Since insulation layers are typically printed using a finer screen, much better edge definition, and thus greater device precision can be achieved. Thus, neither the area of conductive element 16 nor of the integrated reagent/blood separation layer are critical to the performance characteristics of the finished test strip. The conductive elements and the integrated reagent/blood separation layer may therefore be applied using techniques which provide less precision than can be employed in other processes.

It will be appreciated by persons skilled in the art that, while both conductive elements must be accessible to electroactive species in a sample disposed in the sample application region, the important function of the insulation mask is to provide an aperture defining the contact region between conductive element 16 and the integrated reagent/blood separation layer 17. Thus, in the limiting case, it is only necessary to form one aperture in the insulation layer. The second conductive element can be exposed along an edge of the insulation layer, or may be located on a facing surface in a folded electrode structure.

The substrate 10 used in making the test strips of the invention can be any non-conducting, dimensionally stable material suitable for insertion into a glucose test meter. Suitable materials include polyester films, for example a 330 micron

polyester film, and other insulating substrate materials such as polyvinyl chloride (PVC) and polycarbonate.

The conductive elements and associated leads and contacts can be formed from essentially any conductive material including silver, Ag/AgCl, gold, or platinum/carbon, and need not all be formed from the same material. The conductive element 16 is preferably formed from conductive carbon. Preferred conductive carbon are ERCON ERC1, ERCON ERC2 and Acheson Carbon Electrode 423. Carbon with these specifications is available from Ercon Inc. (Waltham, Massachusetts, USA), or Acheson Colloids, (Princes Rock, Plymouth, England). The conductive element 16 makes contact with working electrode track 15, and is close to, but not contacting conductive element 14' disposed as the end of reference electrode track 14.

The insulating layer 18 can be formed from polyester-based printable dielectric materials such as ERCON R488-B(HV)-B2 Blue. The top cover 23 is suitably formed from a polyester strip or a "hot melt" coated plastic.

The test strips of the present invention do not require the formation of a discrete exit port to permit air to escape from the device as sample enters the electrode chamber but instead uses a distributed exit along all of the edges of the mesh. As the sample fluid wicks along the mesh, air seeps out of the edges of the mesh all around the device underneath the top layer. The sample fluid does not seep out because the insulation layer imparts significant hydrophobicity to that part of the mesh. The liquid sample therefore remains in the central hydrophilic region.

The key to the performance achieved using the present invention is in the nature of the integrated reagent/blood separation layer 17. This layer can be formed from a mixture containing a filler which has both hydrophobic and hydrophilic surface regions, and in the case of a glucose test strip, an enzyme which can oxidize glucose, and a mediator which can transfer electrons from the enzyme to the underlying conductive element layer 16. This layer is suitably formed by formulating an ink which contains the filler, the enzyme and the mediator in a suitable carrier and using this ink to print the layer 17 onto the device.

5 A preferred filler for use in the layer 17 is silica. Silica is available in a variety of grades and with a variety of surface modifications. While all silica compounds tested resulted in a product which could measure glucose under some conditions, the superior performance characteristics of glucose test strip of the invention are obtained when a silica having a surface modification to render it partially hydrophobic is used. Materials of this type include Cab-O-Sil TS610, a silica which is modified by partial surface treatment with methyl dichlorosilane; Cab-o-Sil 530, a silica which is modified by full surface treatment with hexamethyl disilazane; Spherisorb C4 silica, which is surface modified with 4 carbon chains; and other similarly modified silicas, or combinations thereof. Silica with a surface modification which is too hydrophobic should be avoided. For example, it has been observed that C18-modified silica is too hydrophobic to form a printable ink.

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15 During the process of manufacturing the ink of the invention, the particles are broken down by homogenization to expose hydrophilic inner portions of the silica particles. The actual particles present in the ink therefore have both hydrophilic and hydrophobic regions. The hydrophilic regions form hydrogen bonds with each other and with water.

20 When this material is formulated into an ink as described below in Example 1, and screen printed onto the conductive element 16, the dual nature of the material causes it to form layers of two-dimensional networks which take form as a kind of honeycomb which is visible upon microscopic examination. On rehydration, this layer does not break up, but swells to form a gelled reaction zone in the vicinity of the underlying conductive element 16. Reactants such as enzyme, mediator and glucose move freely within this zone, but interfering species such as red blood cells containing oxygenated hemoglobin are excluded. This results in a device in which the amount of current generated in response to a given amount of glucose varies by less than 10 percent over a hematocrit range of 40 to 60 %, and which is thus substantially insensitive to the hematocrit of the sample, and in fact performs substantially the same in blood as in a cell-free control solution. (Figs. 3A-C, Fig. 4 and Fig. 5A - 5C)

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Furthermore, the gelled reaction zone presents a greater barrier to entry of blood analytes such as glucose which makes the device diffusion, rather than kinetically limited. This leads to a device in which the measured current varies by less than 10 percent over a temperature range from 20°C to 37°C and which is thus essentially temperature independent. (Figs. 6A and 6B)

When making a glucose test strip, the integrated reagent/blood separation layer is advantageously formed from an aqueous composition containing 2 to 10 % by weight, preferably 4 to 10 % and more preferably about 4.5 % of a binder such as hydroxyethylcellulose or mixtures of hydroxyethylcellulose with alginate or other thickeners; 3 to 10 % by weight, preferably 3 to 5 % and more preferably about 4 % silica; 8 to 20 % by weight, preferably 14 to 18 % and more preferably about 16 % of a mediator such as ferricyanide; and .4 to 2 % by weight, preferably 1 to 2 % and more preferably about 1.6 % of an enzyme such as glucose oxidase, assuming a specific activity of about 250 units/mg, or about 1000 to 5000 units per gram of ink formulation.

The integrated reagent/blood separation layer may also include additional ingredients without departing from the scope of the invention. For example, the nonconducting layer may include an antifoam. In addition, the nonconducting layer may be formulated with a buffering agent to control the pH of the reaction zone. The pH may be maintained at a level within the range from about pH 3 to pH 10. In one embodiment of the invention, it is of particular utility to maintain the pH of the device at a level above 8 because at this pH oxygen bound to hemoglobin is not released. Further, at this pH, the reaction rate of glucose oxidase with oxygen is very low. Thus, selection of an appropriate pH can further stabilize the performance of the test strip against the effects of varying hematocrit. In an alternative embodiment of the invention, maintaining a low pH (below pH 5.5, the optimum pH for reaction of glucose oxidase with oxygen) may be preferred. For example, maintaining a pH of around pH 5 is better if the primary concern is the elimination of electrochemical interferences arising from oxidation of interfering substances such as ascorbic acid, uric acid or acetaminophen, since these compounds are more difficult to oxidize at lower pH.

While a preferred embodiment of the invention is a glucose test strip as described above, the test strips of the invention are not limited to the detection of glucose. For example, a fructosamine test strip could include two layers disposed over the conductive element. The first, lower layer is formed from an ink comprising a carbonate buffer (pH>10) in a silica mix substantially as described in Example 7 but without enzyme, mediator or citrate buffer. The second, upper layer is formed from an ink further comprising an oxidant such as ferricyanide.

Fig. 7 shows an alternative embodiment of the invention. In this embodiment, a second non-conductive layer 71 is disposed over the integrated reagent/blood separation layer 17. This layer is formed from a composition which is identical to the first integrated reagent/blood separation layer except that the enzyme or both the enzyme and the mediator are omitted. This layer further isolates the conductive element 16 from contact with oxygen-carrying red blood cells, thus reducing the effects of oxygen. Furthermore, to the extent that enzyme may tend to diffuse away from the surface of the electrode during the course of the measurement, such a layer containing mediator can provide an increased region in which it will have mediator available for the transfer of electrons.

EXAMPLE 1

A non-conducting formulation for preparation of the integrated reagent/blood separation layer 17 was made as follows. 100 ml of 20 mM aqueous trisodium citrate was adjusted to pH 6 by the addition of 0.1 M citric acid. To this 6 g of hydroxyethyl cellulose (HEC) was added and mixed by homogenization. The mixture was allowed to stand overnight to allow air bubbles to disperse and then used as a stock solution for the formulation of the coating composition.

2 grams Cab-o-Sil TS610 silica and 0.1 grams of Dow Corning antifoam compound was gradually added by hand to 50 grams of the HEC solution until about 4/5 of the total amount had been added. The remainder was added with mixing by homogenization. The mixture was then cooled for ten minutes in a refrigerator. 8 g of potassium hexacyanoferrate (III) was then added and mixed until completely dissolved. Finally, 0.8 g of glucose oxidase enzyme preparation (250

Units/mg) was added and then thoroughly mixed into the solution. The resulting formulation was ready for printing, or could be stored with refrigeration.

EXAMPLE 2

5 To prepare glucose test strips using the ink formulation of Example 1, a series of patterns are used to screen print layers onto a 330 micron polyester substrate (Melinex 329). The first step is the printing of carbon pads. An array of 10 X 50 pads of carbon is formed on the surface of the polyester substrate by printing with EC2 carbon. (Ercon) The printed substrate is then passed through a heated dryer, and optionally cured at elevated temperature (e.g. 70°C) for a period of 1 to 3 weeks.

10 Next, an array of silver/silver chloride connecting tracks and contacts is printed onto the substrate using ERCON R-414 (DPM-68) 1.25 bioelectrode sensor coating material and dried. One working track which makes contact with the carbon pad and one reference track is printed for each carbon pad in the array.

15 A dielectric layer is then printed using ERCON R488-B(HV)-B2 Blue and dried. The dielectric layer is printed in a pattern which covers substantially all of each device, leaving only the contacts, the tip of the reference electrode and the carbon pads uncovered.

20 On top of the dielectric layer the ink of Example 1 is used to form a integrated reagent/blood separation layer overlaid on top of each conductive carbon pad.

25 Polyester mesh strips (Scrynel PET230 HC) are then laid down across the substrate in lines, covering the reactions areas exposed by the windows in the dielectric. An 5 mm wide polyester strip (50 microns thick) is then applied over the top of the mesh strips, and the edges of the electrodes are heat sealed. Finally, the substrate is cut up to provide 50 individual electrodes, for example having a size of 5.5 mm wide and 30 mm long.

EXAMPLE 3

30 Test strips manufactured using the ink formulation of Example 1 in the manner described in Example 2 were placed in a test meter with an applied voltage of

500 mV and used to test blood samples having varying glucose concentrations and hematocrits ranging from 40% to 60%. Figs. 3A-3C show the current measured 25 seconds after applying the voltage as a function of the glucose concentration, and Fig. 4 plots the slope of the glucose response as a function of hematocrit. As can be seen, the indicators produce highly reproducible current levels which are essentially independent of hematocrit.

EXAMPLE 4

Glucose test strips in accordance with the invention were made in accordance with Example 2, except the non-conductive layer was formed with 7 g Spherisorb C4 and 1 g Cab-o-Sil TS610. This formulation was laid down on three different types of carbon-containing conductive elements as follows:

A: Ercon EC1

B: Ercon EC2

C: Ercon EC2 on top of Acheson Carbon, Electrodag 423 SS.

These test strips were used to measure varying levels of glucose in either a control solution (One Touch Control Solution, Lifescan Inc.) containing glucose in an inert solution or in blood at an applied voltage of 425 mV. The current observed 25 seconds after the voltage was applied was measured. Figs. 5A - 5C show the results obtained for the three formulations, A, B, and C, respectively. In all cases, the slope of the line showing the response of the meter to different glucose concentrations was essentially the same whether the measurements were made in blood or the control solution. Thus, this further demonstrates the independence of the test strips of the invention from the oxygen content and hematocrit of the sample, as well as the ability to use various materials as the conductive element.

EXAMPLE 5

Test strips prepared in accordance with Example 2 were tested at two different sample temperatures, namely 37 °C and 20 °C using an applied voltage of 425 mV. Figs. 6A and 6B show the current measured 25 seconds after applying the voltage as a function of glucose concentration. As can be seen, the slopes of the two

lines are essentially identical (0.1068 at 20 °C versus 0.1009 at 37 °C), thus demonstrating that the test strips provide essentially temperature-independent behavior over a temperature range from ambient to physiological temperatures.

EXAMPLE 6

The current transient was measured for a test strip prepared in accordance with Example 2 and for a commercial test strip made with a carbon-containing ink. The results are shown in Figs. 8A and 8B. As shown, the test strip of the invention (Fig. 8A) provides a very flat transient which maintains more than 50% of the peak current for a period of more than 25 seconds after the initial response from the test strip. In contrast, the carbon-based electrode exhibited an almost immediate decay in the current, having lost 50% of the peak current in a period of the first 1 to 2 seconds after the initial response from the test strip. This makes timing of the measurement difficult if peak current values are to be captured, or reduces the dynamic range of the meter if current must be measured after substantial decay has occurred. Thus, the test strips of the invention are advantageous in that the current generated in response to a given amount of glucose decays by less than 50% in the 5 seconds following peak current generation.

EXAMPLE 7

An ink for printing glucose test strips in accordance with the invention was formulated as follows:

67.8 g 20 mM Citrate buffer, pH 6
0.68 g Polyvinyl alcohol (MW 85,000-146,000, 88% hydrolysed)
0.68 g of Polyvinyl pyrrolidone-vinyl acetate
0.42 g of Dow Corning DC1500 antifoam
3.4 g of hydroxyethyl cellulose (Natrosol 250G, Hercules)
5.5 g of surface modified silica (Cabo-Sil TS 610, Cabot)
1.5 g glucose oxidase
20.0 g Potassium Ferricyanide.

EXAMPLE 8

Figs. 10A-I shows the stepwise preparation of a test strip in accordance with the invention. As is apparent from a comparison of this test strip and the strip of Fig. 1, the precise arrangement of the electrodes on the strip is not critical. Further, different materials may be used in fabricating the strip.

The first step in fabricating the test strip is the deposition of silver tracks 101, 102 of substrate 100. A preferred substrate is a 500 micron thick polyester film sold under the tradename Valox™. The silver electrodes can be formed by screen printing using an ink composition formulated as in Example 2.

After deposition of the silver electrodes, a second electrode print is carried out to form carbon conductive elements 103, 104 and 105 as shown in Fig 10B. Conductive element 103 is formed in contact with silver track 101 and will form the working electrode in the finished test strip. Carbon pads 104 and 105 connect electrically to the ends of silver tracks 101 and 102 and provide connection between the strip and a test meter. The carbon conductive elements can be formed by screen printing with a conductive carbon ink formulation such as those described in the previous examples.

The next step in the manufacturing process is the deposition of an insulation layer 106 for example by screen printing an insulation ink, for example the dielectric ink of Example 2. (Fig. 10C) As shown, the insulation layer contains three windows 107, 108 and 109. Window 108 is aligned with the end of the carbon conductive element 103. Window 107 is aligned with the end of silver track 102 to provide access to the reference electrode. The third window, 109, is provided to permit passage of insulation material from the second insulation coating through the mesh layer, but is not required.

Fig 10D shows the next step in the process, which is the formation of an integrated reagent/blood separation layer 110. This layer is deposited over window 108 and extends over the insulation layer 106 along all sides of the window 108. A suitable formulation for printing layer 110 has the following composition to provide an integrated reagent/blood separation layer with a buffered pH of about 6:

Component	Amount
Analar Water	3L
Tri-sodium Citrate	15.75g
Nat 250 G	150g
Citric Acid	6.3g
Poly Vinyl Alcohol	30g
DC 1500 Defoamer	15 ml
Cabosil	225 g
Glucose Oxidase	48g
Potassium Hex/60299	660g
PVPVA	30g

After the integrated reagent/blood separation layer 110 is formed, a layer of mesh 111 is deposit over the sample collection region of the test strip. (Fig. 10E) The mesh 111 is preferably a nylon mesh which has been pretreated with acetone and Fluorad FC 170C surfactant to render the mesh hydrophilic. The purpose of the mesh 111 is the transport of the liquid sample evenly through the area between the working and reference electrodes.

A second insulation print 112 is then carried out using a slightly more flexible insulation ink (ERCON Insulayer 820202) to define the sample collection region. (Fig. 10F). A tape cover 113 is then applied over the top of the test strip as described above in Example 2 to form a finished test strip. (Fig. 10G).

CLAIMS

1 1. A disposable test strip for use in a test meter of the type which
2 receives a disposable test strip and a sample of blood and performs an electrochemical
3 analysis of the amount of a blood analyte in the sample, comprising:

4 (a) a substrate;
5 (b) a first conductive element disposed on the substrate;
6 (c) a second conductive element disposed on the substrate in
7 sufficient proximity to the first conductive element to allow the completion of an
8 electrical circuit between the first and second conductive elements when a sample of
9 blood is placed on the test strip;

10 (d) a non-conductive integrated reagent/blood separation layer
11 disposed over the first conductive element, said integrated reagent/blood separation
12 layer comprising reagents for the electrochemical detection of the analyte dispersed in
13 a non-conductive matrix effective to exclude blood cells from the surface of the first
14 conductive element while permitting access to the first conductive element by soluble
15 electroactive species;

16 (e) contacts for making an electrical connection between the first
17 and second conductive elements and the test meter; and

18 (f) an insulation layer disposed over at least the first conductive
19 elements, said insulation layer having a first aperture therein aligned with the first
20 conductive element, wherein the non-conductive integrated reagent/blood separation
21 layer contacts the first conductive element through the aperture in the insulation layer
22 and wherein the non-conductive integrated reagent/blood separation layer is formed
23 covering the entire first aperture, thereby leaving no portion of the first conductive
24 element directly exposed to a sample applied to the test strip.

1 2. The test strip of claim 1, wherein the integrated reagent/blood
2 separation layer comprises an enzyme for oxidation of glucose and a redox mediator
3 effective to transfer electrons from the enzyme to the first conductive element.

1 3. The test strip of claim 2, wherein the matrix comprises silica
2 having both hydrophobic and hydrophilic surfaces.

1 4. The test strip of claims 3, wherein the first and second
2 conductive elements comprise conductive carbon.

1 5. The test strip according to claim 4, wherein the enzyme is
2 glucose oxidase.

1 6. The test strip according to claim 5, wherein the redox mediator
2 is ferricyanide.

1 7. The test strip according to claim 3, wherein the integrated
2 reagent/blood separation layer is formed from an aqueous composition comprising 2
3 to 10 % by weight of a binder; 3 to 10 % by weight of silica; 8 to 20 % by weight of
4 the redox mediator; and 1000 to 5000 units of the enzyme per gram of the aqueous
5 composition.

1 8. The test strip of claims 7, wherein the first and second
2 conductive elements comprise conductive carbon.

1 9. The test strip according to claim 7, wherein the enzyme is
2 glucose oxidase.

1 10. The test strip according to claim 9, wherein the redox mediator
2 is ferricyanide.

1 11. A method for forming a disposable test strip for use in a test
2 meter of the type which receives a disposable test strip and a sample of blood and
3 performs an electrochemical analysis of the amount of a blood analyte in the sample,
4 comprising:

5 (a) forming first and second conductive elements on a substrate;
6 (b) forming a layer of insulation covering the first conductive
7 element, said layer of insulation having a first aperture therein aligned with a portion
8 of the first conductive element in a sample application region; and
9 (c) forming a reagent layer disposed on the insulation layer and
10 making contact with the first conductive element through the first aperture in the
11 insulation layer, said integrated reagent/blood separation layer comprising reagents for
12 the electrochemical detection of glucose dispersed in a non-conductive matrix
13 effective to exclude blood cells from the surface of the first conductive element while
14 permitting access to the first conductive species by soluble electroactive species,
15 whereby the first conductive element is isolated from direct contact with a sample
16 placed on the test strip.

1 12. The method of claim 11, wherein the reagent layer is a non-
2 conductive integrated reagent/blood separation layer.

1 13. The method of claim 12, wherein the non-conductive
2 integrated reagent/blood separation layer is formed covering the entire first aperture,
3 thereby leaving no portion of the first conductive element directly exposed to a
4 sample applied to the test strip.

1 14. The method of claim 13, wherein the integrated reagent/blood
2 separation layer comprises an enzyme for oxidation of glucose and a redox mediator
3 effective to transfer electrons from the enzyme to the first conductive element.

1 15. The method of claim 14, wherein the matrix is formed from
2 silica having both hydrophobic and hydrophilic surfaces.

1 16. The method of claims 15, wherein the first and second
2 conductive elements comprise conductive carbon.

1 17. The method of claim 16, wherein the enzyme is glucose
2 oxidase.

1 18. The method of to claim 17, wherein the redox mediator is
2 ferricyanide.

1 19. The method of claim 13, wherein the integrated reagent/blood
2 separation layer is formed from an aqueous composition comprising 2 to 10 % by
3 weight of a binder; 3 to 10 % by weight of silica; 8 to 20 % by weight of the redox
4 mediator; and 1000 to 5000 units of the enzyme per gram of the aqueous composition.

1 20. The method of claims 19, wherein the first and second
2 conductive elements comprise conductive carbon.

1 21. The method of claim 19, wherein the enzyme is glucose
2 oxidase.

1 22. The method of claim 21, wherein the redox mediator is
2 ferricyanide.

1 23. The method of claim 13, wherein the insulation layer is formed
2 over both the first and second conductive elements, and includes a second aperture
3 aligned with the second conductive element.

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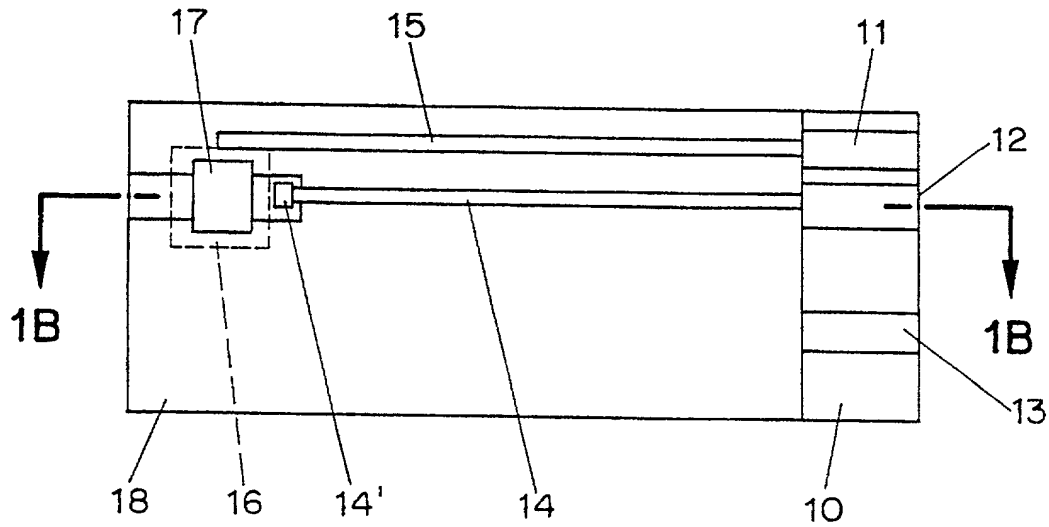


FIG. 1A

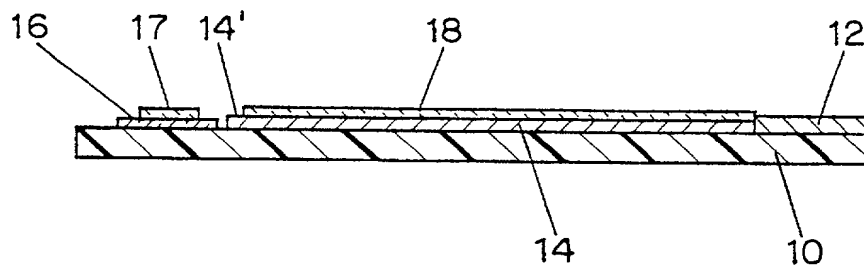


FIG. 1B

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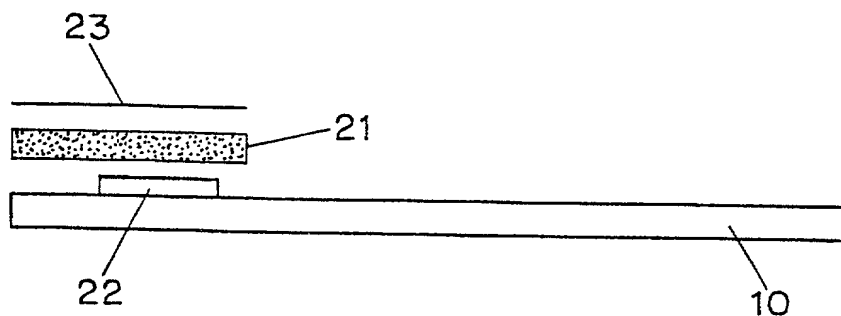


FIG. 2

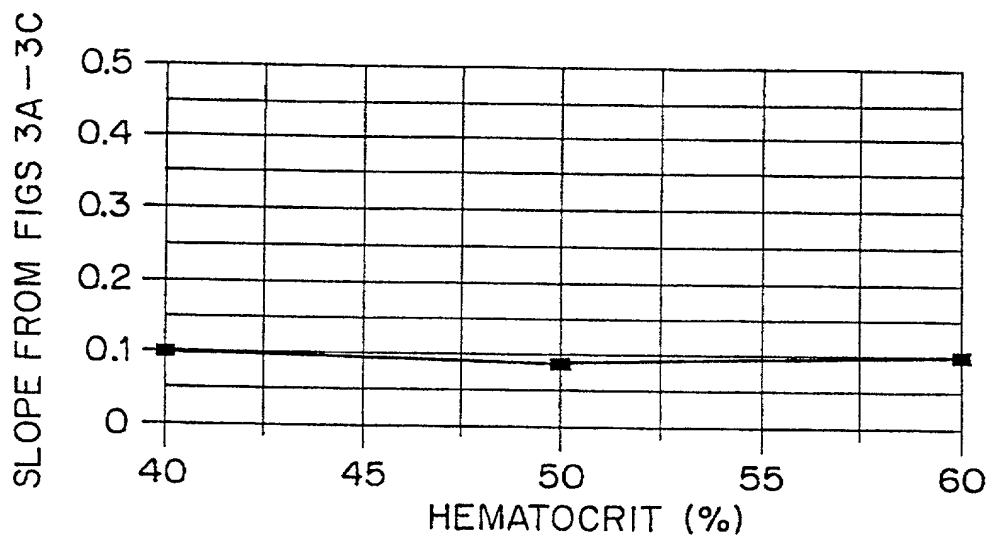


FIG. 4

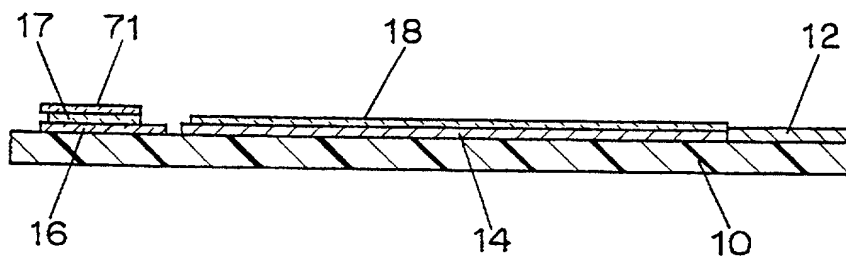


FIG. 7

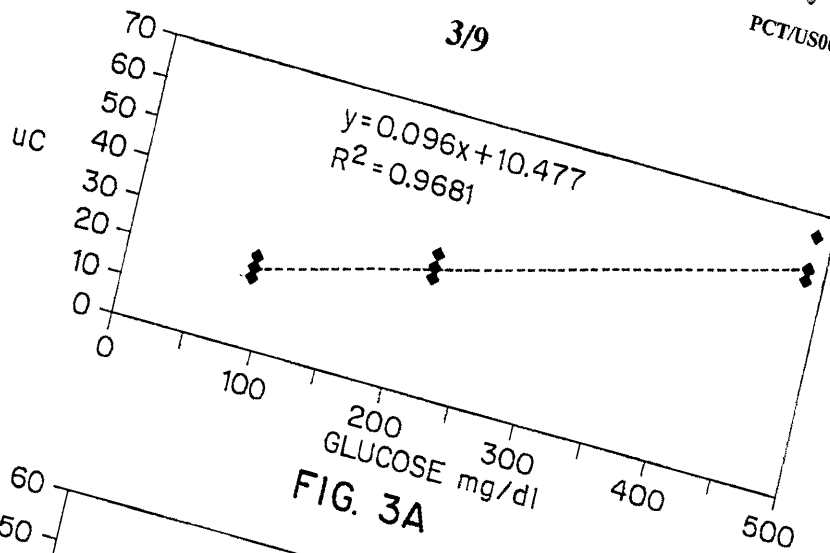


FIG. 3A

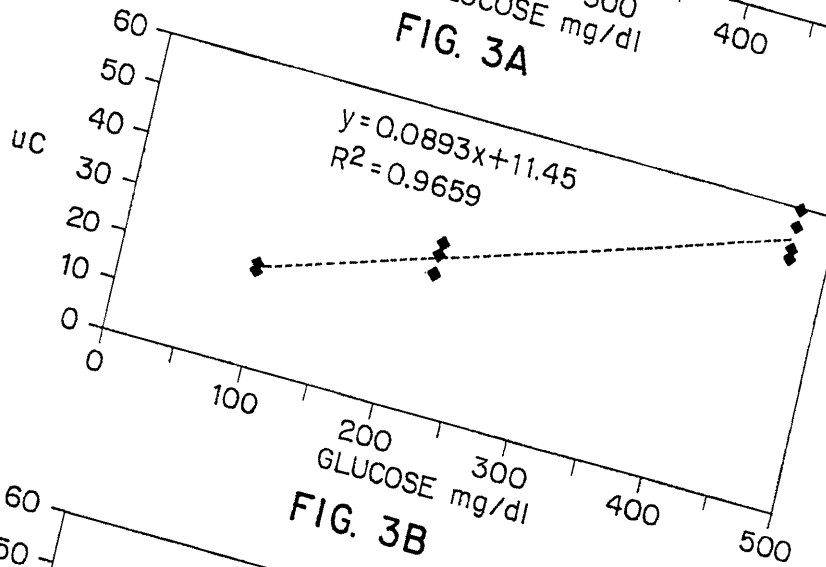


FIG. 3B

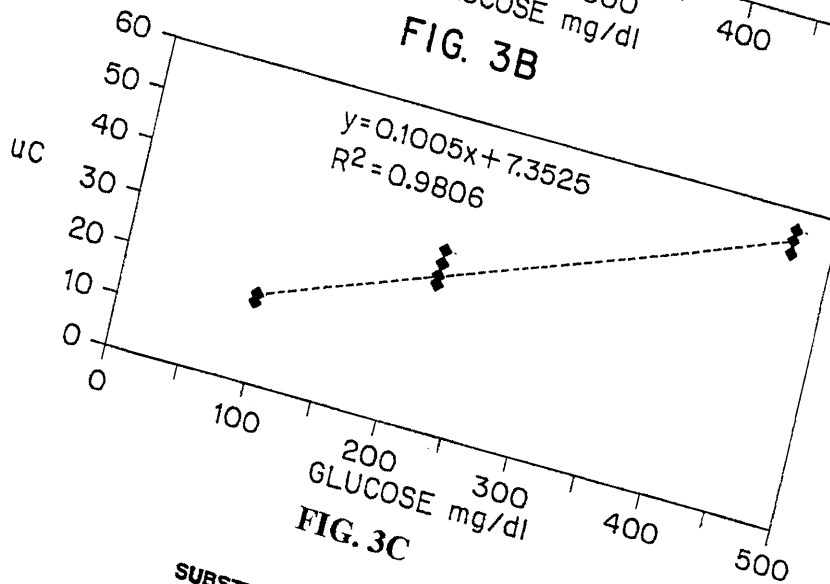
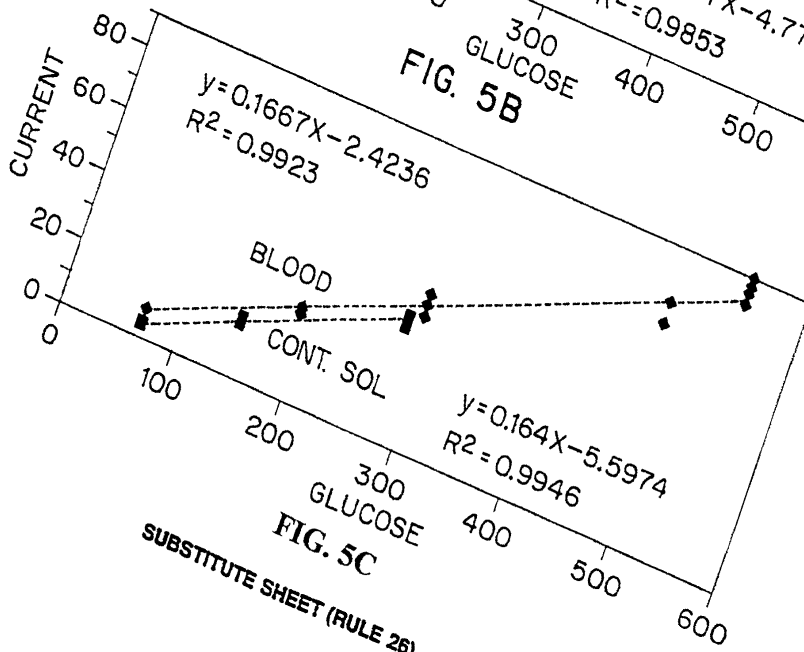
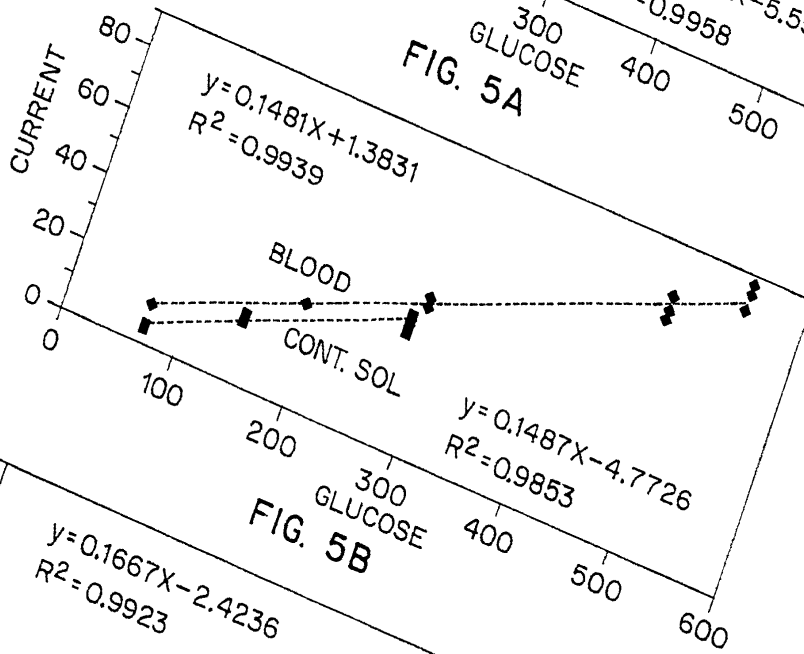
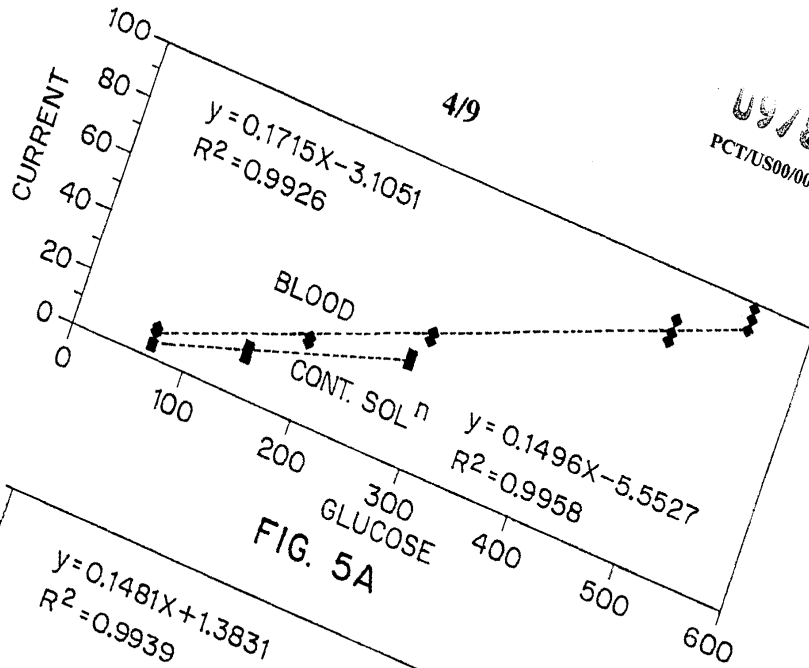


FIG. 3C



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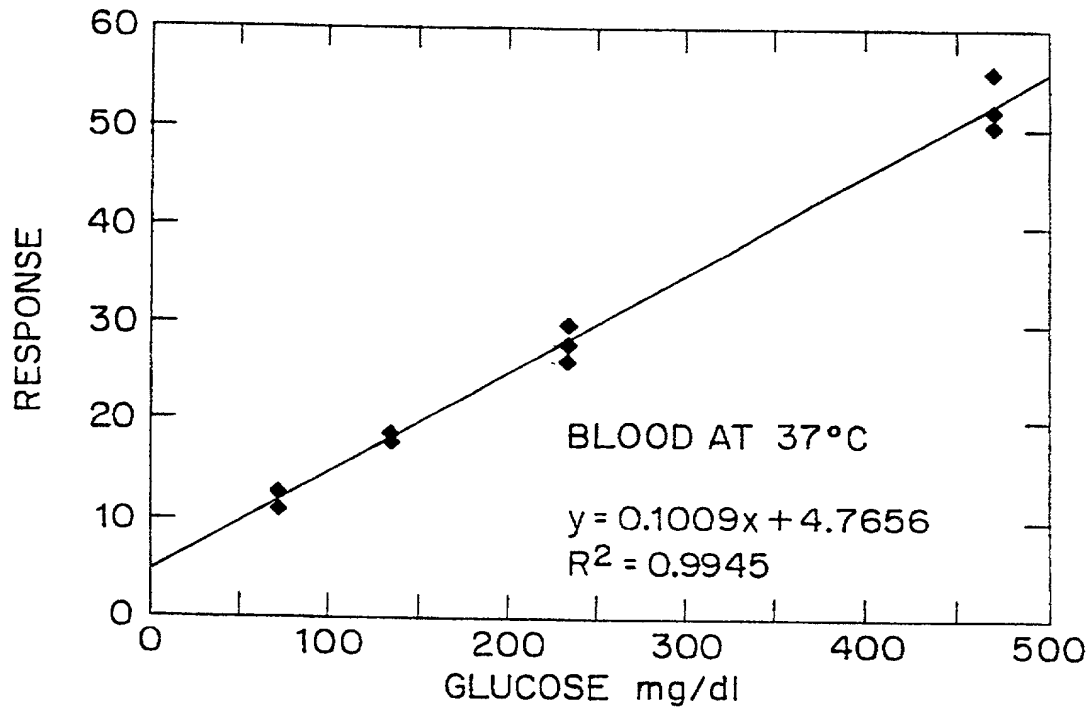


FIG. 6A

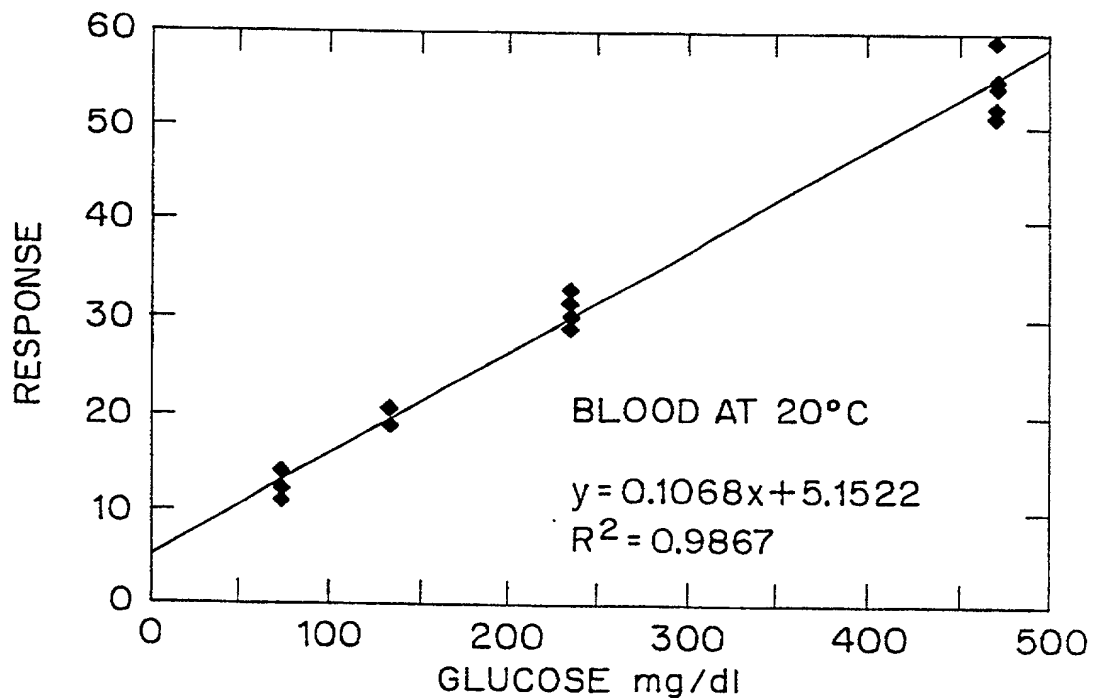


FIG. 6B

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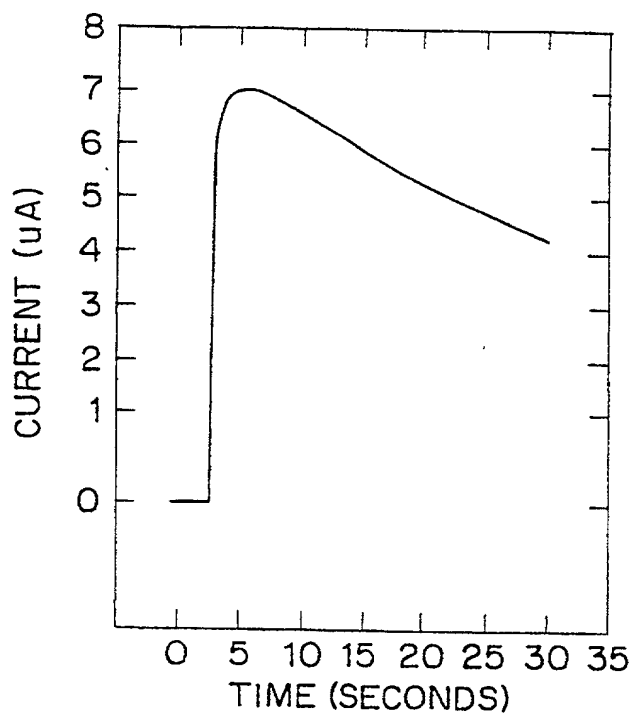


FIG. 8A

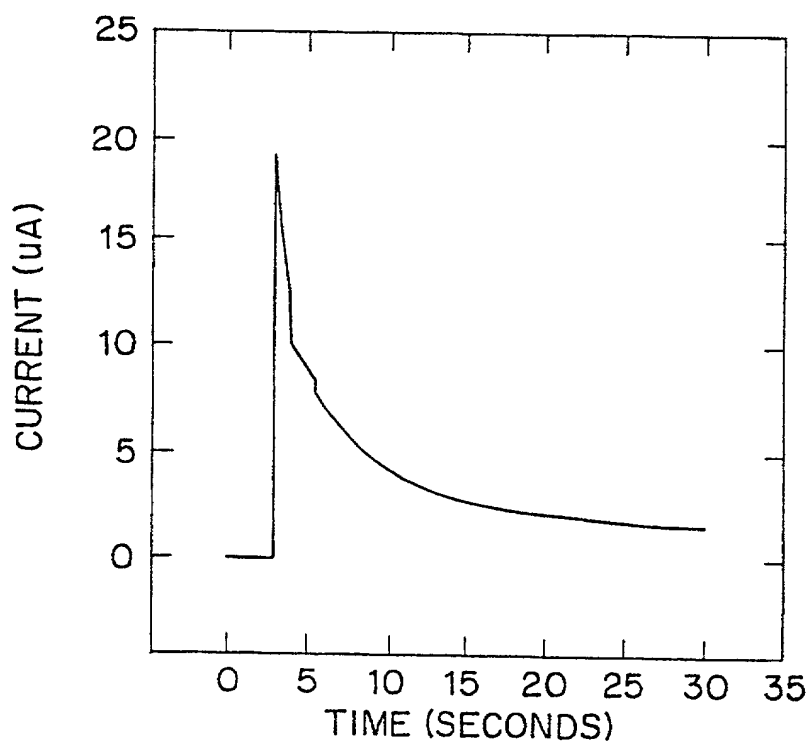


FIG. 8B

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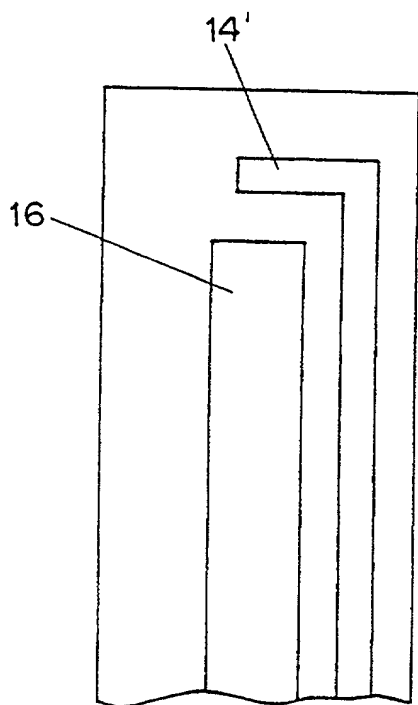


FIG. 9A

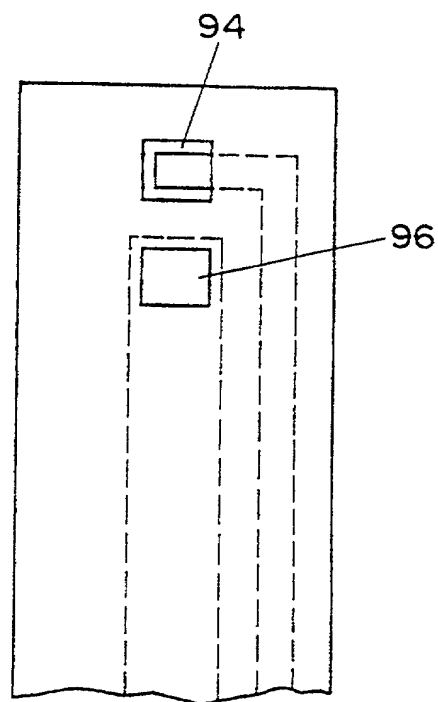


FIG. 9B

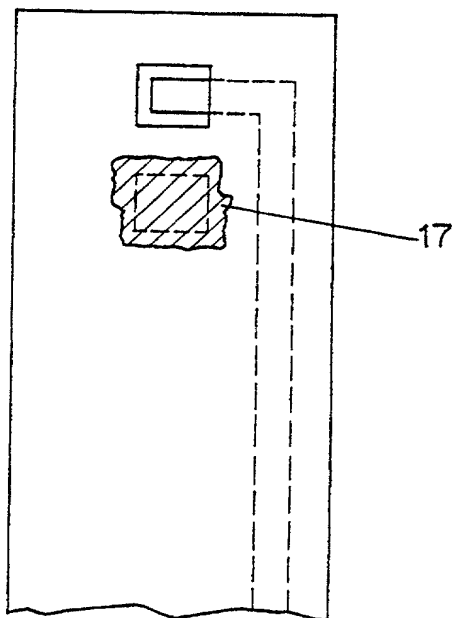


FIG. 9C

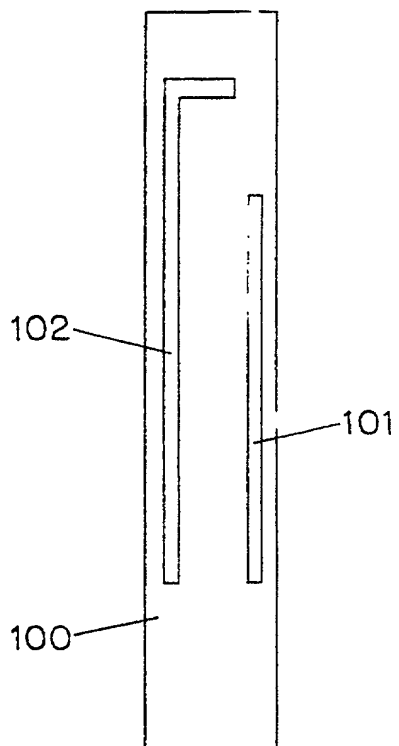


FIG. 10A

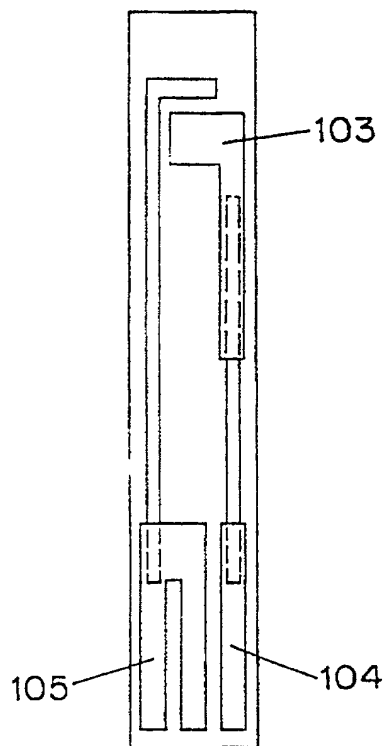


FIG. 10B

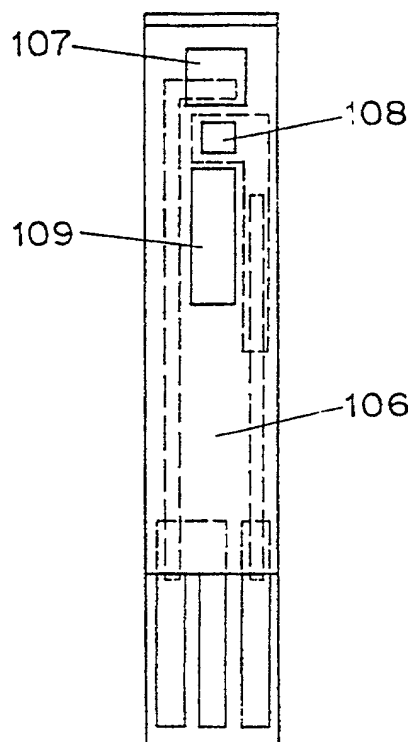


FIG. 10C

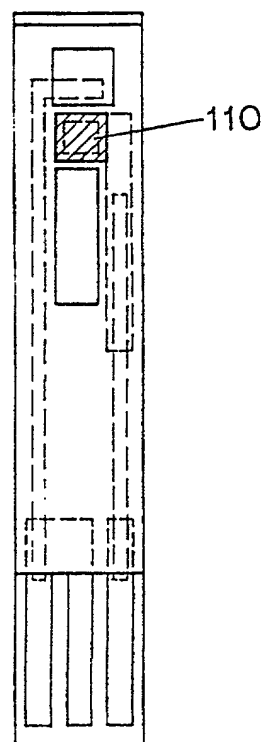


FIG 10D

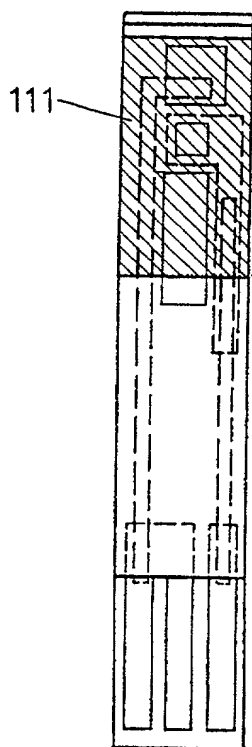


FIG. 10E

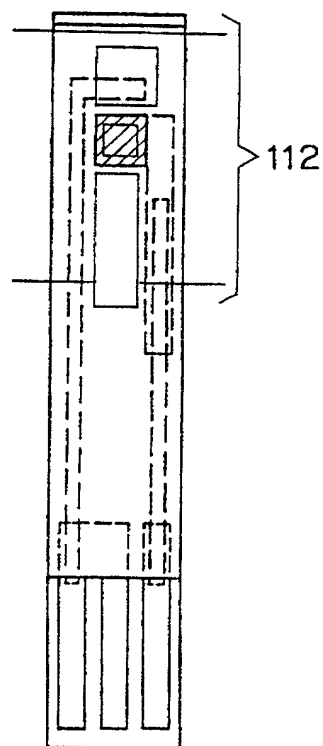


FIG. 10F

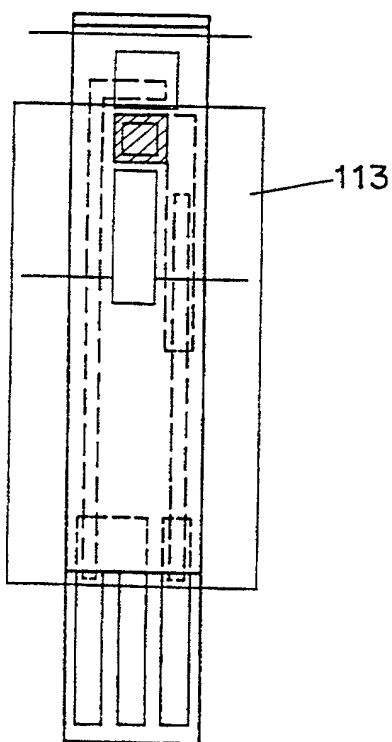


FIG. 10G

Claim for Priority

I hereby claim foreign priority benefits under 35 U.S.C. § 119 (a)-(d) or 365(b) of any foreign application(s) for patent or inventor's certificate, or 365(a) of any PCT International application which designated at least one country other than the United States of America, listed below and have also identified below any foreign applications for patent or inventor's certificate, or of any PCT International application having a filing date before that of the application on which priority is claimed.

EARLIEST FOREIGN APPLICATION(S). FILED WITHIN TWELVE MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION					
COUNTRY	APPLICATION NO.	DATE OF FILING (day-month-year)	DATE OF ISSUE (day-month-year)	PRIORITY CLAIMED	CERTIFIED COPY ATTACHED
				YES [] NO []	YES [] NO []
FOREIGN APPLICATION(S). IF ANY. FILED MORE THAN 12 MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION					
COUNTRY	APPLICATION NO.	DATE OF FILING (day-month-year)	DATE OF ISSUE (day-month-year)		


Provisional Application

I hereby claim the benefit under 35 U.S.C. § 119(e) of any United States provisional application(s) listed below.

(application number)

(filing date)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

NAME OF SOLE OR FIRST INVENTOR	LAST NAME <u>MCALDER</u>	FIRST NAME <u>JEROME</u>	MIDDLE NAME <u>F.</u>
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE <u>Wantage</u>	STATE OR COUNTRY OF RESIDENCE <u>UK G.B.X.</u>	COUNTRY OF CITIZENSHIP <u>UK</u>
POST OFFICE ADDRESS <u>52 Nobles Close</u>		CITY <u>WANTAGE</u>	STATE/COUNTRY ZIP CODE <u>UK OX12 0NR</u>
DATE <u>13-02-02</u>		SIGNATURE 	

- (X) Signature for additional joint inventor attached. Number of Pages 1
() Signature by Administrator (Att) or legal representative for deceased or incapacitated inventor. Number of Pages
() Signature for inventor who refuses to sign or cannot be reached by person authorized under 37 CFR § 1.47. Number of Pages

T-357 P.04/10 F-123

T-058 P.04/04 F-268

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DATE		SIGNATURE	
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POST OFFICE ADDRESS 58 Newland Mill		CITY WITNEY OXON	STATE/COUNTRY ZIP CODE UK OX8 8SZ
DATE		SIGNATURE	
NAME OF SIXTH INVENTOR	LAST NAME DAVIES	FIRST NAME OLIVER	MIDDLE NAME W.F.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 16 Drummond Circle		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 4QP
DATE		SIGNATURE	

**COMBINED DECLARATION
AND POWER OF ATTORNEY**

As a below named inventor, I hereby declare that:

My citizenship, residence and post office address are as listed below next to my name.

I believe I am the original, first and ☐ sole/☒ joint inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled: **Disposable Test Strips With Integrated Reagent/Blood Separation Layer**

the specification of which

(a) ☐ is attached hereto.

(b) ☒ was filed on July 5, 2001 as Application Serial No. 09/869,887 and was amended on _____.

(c) ☐ was described and claimed in International Application No. PCT/US00/00620 filed on January 11, 2001 and amended on _____.

Acknowledgment of Duty of Disclosure

I hereby state that I have reviewed and understood the content of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to the patentability of the subject matter claimed in this application in accordance with Title 37, Code of Federal Regulations § 1.56(a).

35 U.S.C. § 120

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) or 365(c) of any PCT international application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT international application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose material information as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application:

<u>09/228,855</u>	<u>01/12/99</u>	<u>Patented</u>	<u>6,241,862</u>
(Application Serial No.)	(Filing Date)	(Status)(patented,pending,abandoned)	(Patent No. if applicable)

_____ (Application Serial No.)	_____ (Filing Date)	_____ (Status)(patented,pending,abandoned)	_____ (Patent No. if applicable)
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Power of Attorney

I hereby appoint Carl Oppedahl, PTO Reg. No. 32,746, Marina T. Larson, PTO Reg. No. 32,038, and D'Arcy Straub, PTO Reg. No. 47,113 of the firm of OPPEDAHL & LARSON LLP, having office at P.O. Box 5068, Dillon, CO 80435-5068 as attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith.

SEND CORRESPONDENCE TO:



021121

DIRECT TELEPHONE CALLS TO:

OPPEDAHL & LARSON LLP
(870)468-8600

Claim for Priority

I hereby claim foreign priority benefits under 35 U.S.C. § 119 (a)-(d) or 365(b) of any foreign application(s) for patent or inventor's certificate, or 365(a) of any PCT international application which designated at least one country other than the United States of America, listed below and have also identified below any foreign applications for patent or inventor's certificate, or of any PCT international application having a filing date before that of the application on which priority is claimed.

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COUNTRY	APPLICATION NO.	DATE OF FILING (day/month/year)	DATE OF ISSUE (day/month/year)	PRIORITY CLAIMED	CERTIFIED COPY ATTACHED
				YES [] NO []	YES [] NO []
FOREIGN APPLICATION(S), IF ANY, FILED MORE THAN 12 MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION					
COUNTRY	APPLICATION NO.	DATE OF FILING (day/month/year)	DATE OF ISSUE (day/month/year)		

Provisional Application

I hereby claim the benefit under 35 U.S.C. § 119(e) of any United States provisional application(s) listed below.

(application number)

(filing date)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

NAME OF SOLE OR FIRST INVENTOR	LAST NAME MCALEER	FIRST NAME JEROME	MIDDLE NAME F.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE Wantage	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 52 Nobles Close		CITY WANTAGE	STATE/COUNTRY ZIP CODE UK OX12 ONR
DATE		SIGNATURE	

- ☒ Signature for additional joint inventor attached. Number of Pages 1
- ☐ Signature by Administrator(trix) or legal representative for deceased or incapacitated inventor. Number of Pages
- ☐ Signature for inventor who refuses to sign or cannot be reached by person authorized under 37 CFR § 1.47. Number of Pages

#4

COMBINED DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My citizenship, residence and post office address are as listed below next to my name.

I believe I am the original, first and ☐ sole/☒ joint inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled: **Disposable Test Strips With Integrated Reagent/Blood Separation Layer**

the specification of which

(a) ☐ is attached hereto.

(b) ☒ was filed on July 5, 2001 as Application Serial No. 09/869,887 and was amended on _____.

(c) ☐ was described and claimed in International Application No. PCT/US00/00620 filed on January 11, 2001 and amended on _____.

Acknowledgment of Duty of Disclosure

I hereby state that I have reviewed and understood the content of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to the patentability of the subject matter claimed in this application in accordance with Title 37, Code of Federal Regulations § 1.56(a).

35 U.S.C. § 120

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) or 365(c) of any PCT International application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose material information as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT International filing date of this application:

Application Serial No.	Filing Date	Status (patented, pending, abandoned)	Patent No. (if applicable)
09/228,865	01/12/88	Patented	5,241,862
(Application Serial No.)	(Filing Date)	(Status) (patented, pending, abandoned)	(Patent No. if applicable)
(Application Serial No.)	(Filing Date)	(Status) (patented, pending, abandoned)	(Patent No. if applicable)

Power of Attorney

I hereby appoint Carl Oppedahl, PTO Reg. No. 32,746, Marina T. Larson, PTO Reg. No. 32,038, and D'Arcy Straub, PTO Reg. No. 47,113 of the firm of OPPEDAHL & LARSON LLP, having office at P.O. Box 6068, Dillon, CO 80435-8068 as attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith.

SEND CORRESPONDENCE TO:



021121

NOT RECORDED

DIRECT TELEPHONE CALLS TO:
OPPEDAHL & LARSON LLP
(870) 468-6600

09/228,865

2-00

NAME OF SECOND INVENTOR	LAST NAME ALVAREZ-ICAZA	FIRST NAME MANUEL	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK GBX	COUNTRY OF CITIZENSHIP UK MEXICO HAI 15 Feb 2002
POST OFFICE ADDRESS 30 Hill Street 2 Hill Park HAI 15 Feb 2002		CITY HAI Inverness 15 Feb 2002	STATE/COUNTRY ZIP CODE UK IV2 4AL HAI 15 Feb 2002
DATE 15 Feb 2002		SIGNATURE 	

3-00

NAME OF THIRD INVENTOR	LAST NAME HALL	FIRST NAME GEOFF	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK GBX	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS Wilsons Camfield Lodge CARSEWOOD 6 Feb		CITY FORROSC INVERNESS 6 Feb	STATE/COUNTRY ZIP CODE UK IV2 3HS 1010 8-571
DATE 15.2.02		SIGNATURE 	

4-00

NAME OF FOURTH INVENTOR	LAST NAME PLOTKIN	FIRST NAME ELLIOT	MIDDLE NAME V.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK GBX	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 25 Broadstone Park		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 3JZ
DATE 15 Feb 2002		SIGNATURE Elliot Plotkin	

NAME OF FIFTH INVENTOR	LAST NAME SCOTT	FIRST NAME DAVID	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE WITNEY OXON	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 68 Newland mill		CITY WITNEY OXON	STATE/COUNTRY ZIP CODE UK OX8 6S2
DATE		SIGNATURE	

6-00

NAME OF SIXTH INVENTOR	LAST NAME DAVIES	FIRST NAME OLIVER	MIDDLE NAME W.H.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK GBX	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 16 Drummond Gt AN (LWARAN) CROY 15 Feb 2002		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 4GP HAI 15 Feb 2002
DATE 15 Feb 2002		SIGNATURE 	

NAME OF SECOND INVENTOR	LAST NAME ALVAREZ-ICAZA	FIRST NAME MANUEL	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 3C Reay Street		CITY UK	STATE/COUNTRY ZIP CODE UK IV2 3AJ
DATE		SIGNATURE	
NAME OF THIRD INVENTOR	LAST NAME HALL	FIRST NAME GEOFF	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS Willboa Damfield Lane		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 3H5
DATE		SIGNATURE	
NAME OF FOURTH INVENTOR	LAST NAME PLOTKIN	FIRST NAME ELLIOT	MIDDLE NAME V.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 25 Broadstone Park		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 3JZ
DATE		SIGNATURE	
NAME OF FIFTH INVENTOR	LAST NAME <u>SCOTT</u>	FIRST NAME DAVID	MIDDLE NAME
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE <u>WITNEY OXON</u>	STATE OR COUNTRY OF RESIDENCE UK <u>GBK</u>	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 68 Newland mill		CITY WITNEY OXON	STATE/COUNTRY ZIP CODE UK OX8 6S2
DATE 12th February 02		SIGNATURE <u>DS</u>	
NAME OF SIXTH INVENTOR	LAST NAME DAVIES	FIRST NAME OLIVER	MIDDLE NAME W.H.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE INVERNESS	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 15 Drummond Circus		CITY INVERNESS	STATE/COUNTRY ZIP CODE UK IV2 4QP
DATE		SIGNATURE	

5000

COMBINED DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My citizenship, residence and post office address are as listed below next to my name.

I believe I am the original, first and ☐ sole/☒ joint inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled: **Disposable Test Strips With Integrated Reagent/Blood Separation Layer**

the specification of which

(a) ☐ is attached hereto.

(b) ☒ was filed on July 5, 2001 as Application Serial No. 09/869,887 and was amended on _____

(c) ☐ was described and claimed in International Application No. PCT/US00/00620 filed on January 11, 2001 and amended on _____

Acknowledgment of Duty of Disclosure

I hereby state that I have reviewed and understood the content of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to the patentability of the subject matter claimed in this application in accordance with Title 37, Code of Federal Regulations § 1.56(a).

35 U.S.C. § 120

I hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s) or 365(c) of any PCT international application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT international application in the manner provided by the first paragraph of 35 U.S.C. § 112, I acknowledge the duty to disclose material information as defined in 37 CFR § 1.56 which became available between the filing date of the prior application and the national or PCT international filing date of this application:

<u>09/228,855</u>	<u>01/12/99</u>	<u>Patented</u>	<u>6,241,862</u>
(Application Serial No.)	(Filing Date)	(Status)(patented,pending,abandoned)	(Patent No. if applicable)
_____ (Application Serial No.)	_____ (Filing Date)	_____ (Status)(patented,pending,abandoned)	_____ (Patent No. if applicable)

Power of Attorney

I hereby appoint Carl Oppedahl, PTO Reg. No. 32,746, Marina T. Larson, PTO Reg. No. 32,038, and D'Arcy Straub, PTO Reg. No. 47,113 of the firm of OPPEDAHL & LARSON LLP, having office at P.O. Box 5068, Dillon, CO 80435-5068 as attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith.

<p>SEND CORRESPONDENCE TO:</p> <p>021121</p>	<p>DIRECT TELEPHONE CALLS TO:</p> <p>OPPEDAHL & LARSON LLP</p> <p>(970)468-6600</p>
---	---

Claim for Priority

I hereby claim foreign priority benefits under 35 U.S.C. § 119 (a)-(d) or 365(b) of any foreign application(s) for patent or inventor's certificate, or 365(a) of any PCT International application which designated at least one country other than the United States of America, listed below and have also identified below any foreign applications for patent or inventor's certificate, or of any PCT International application having a filing date before that of the application on which priority is claimed.

EARLIEST FOREIGN APPLICATION(S), FILED WITHIN TWELVE MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION					
COUNTRY	APPLICATION NO.	DATE OF FILING (day/month/year)	DATE OF ISSUE (day/month/year)	PRIORITY CLAIMED	CERTIFIED COPY ATTACHED
				YES [] NO []	YES [] NO []

FOREIGN APPLICATION(S), IF ANY, FILED MORE THAN 12 MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION				
COUNTRY	APPLICATION NO.	DATE OF FILING (day/month/year)	DATE OF ISSUE (day/month/year)	

Provisional Application

I hereby claim the benefit under 35 U.S.C. § 119(e) of any United States provisional application(s) listed below.

(application number)

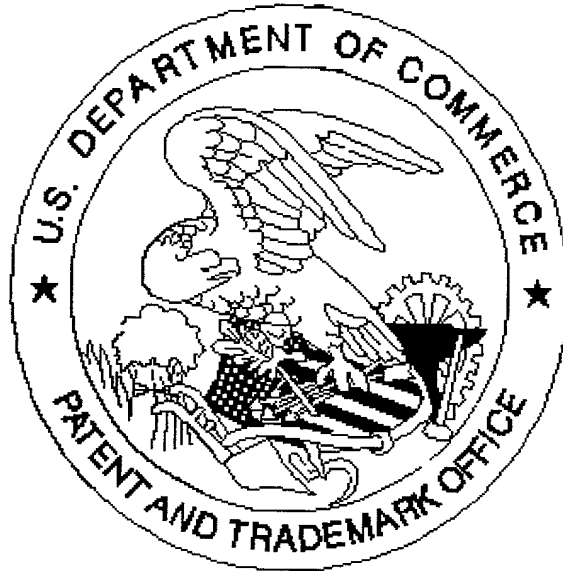
(filing date)

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

NAME OF SOLE OR FIRST INVENTOR	LAST NAME MCALEER	FIRST NAME JEROME	MIDDLE NAME F.
RESIDENCE & CITIZENSHIP	CITY OF RESIDENCE Wantage	STATE OR COUNTRY OF RESIDENCE UK	COUNTRY OF CITIZENSHIP UK
POST OFFICE ADDRESS 52 Nobles Close		CITY WANTAGE	STATE/COUNTRY ZIP CODE UK OX12 ONR
DATE		SIGNATURE	

- [X] Signature for additional joint inventor attached. Number of Pages 1
 [] Signature by Administrator (pri) or legal representative for deceased or incapacitated inventor. Number of Pages ____
 [] Signature for inventor who refuses to sign or cannot be reached by person authorized under 37 CFR § 1.47. Number of Pages ____

United States Patent & Trademark Office
Office of Initial Patent Examination -- Scanning Division



Application deficiencies found during scanning:

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for scanning. (Document title)

☐ Page(s) _____ of _____ were not present
for scanning. (Document title)

☒ *Scanned copy is best available. declaration is dark*